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**INFORMATION
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U.S. PATENT DOCUMENTS

Examiner Initial	Document Number	Date	Name	Class	SubClass	Filing Date If Appropriate
MPA	5,741,645	04/21/98	Orr et al.			

FOREIGN PATENT DOCUMENTS

		Document Number	Date	Country	Class	SubClass	Translation	
							Yes	No

OTHER DOCUMENTS (Including Authors, Title, Date, Pertinent Papers, etc.)

MPA	/	Banfi, et al., "Identification and characterization of the gene causing type 1 spinocerebellar ataxia," <u>Nature Genetics</u> , <u>7</u> , 513-519 (1994).
MPA	/	Filla et al., "Prevalence of hereditary ataxias and spastic paraplegias in Molise, a region of Italy," <u>J. Neurol.</u> , <u>239</u> , 351-353 (1992).
MPA	/	Kremer, et al., "Mapping of DNA Instability at the Fragile X to a Trinucleotide Repeat Sequence p(CCG)n," <u>Science</u> , <u>252</u> , 1711-1714 (1991).
MPA	/	MacDonald et al., "A Novel Gene Containing a Trinucleotide Repeat That Is Expanded and Unstable on Huntington's Disease Chromosomes," <u>Cell</u> , <u>72</u> , 971-983 (1993).
MPA	/	Mahadevan, et al., "Myotonic Dystrophy Mutation: An Unstable CTG Repeat in the 3' Untranslated Region of the Gene," <u>Science</u> , <u>255</u> , 1253-1255 (1992).
MPA	/	Polo et al., "Hereditary Ataxias and Paraplegias in Cantabria, Spain," <u>Brain</u> , <u>114</u> , 855-866 (1991). ^{Hereditary}
MPA	/	Rubensztein, et al., Phenotypic Characterization of Individuals with 30-40 CAG Repeats in the Huntington Disease (HD) Gene Reveals HD Cases with 36 Repeats and Apparently Normal Elderly Individuals with 36-39 Repeats," <u>Am. J. Hum. Genet.</u> , <u>59</u> , 16-22 (1996).

EXAMINER*mpalle***Date Considered**

6/10/99

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